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4) Title: CONTRAST AGENT	IS FOR DIAGNOSTIC	IMAGIN	G .
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CONTRAST AGENTS FOR DIAGNOSTIC IMAGING TECHNICAL FIELD OF THE INVENTION

The present invention relates to diagnostic

nuclear magnetic resonance imaging (MRI or MR imaging).

In particular, this invention relates to novel
compounds that upon administration to a patient, can
enhance the contrast in images of the patient's organs
and tissue obtained through MRI. This invention also
relates to pharmaceutical compositions comprising these
compounds and to methods of making and using the
compounds and compositions for MRI contrast
enhancement.

BACKGROUND OF THE INVENTION

Nuclear magnetic resonance imaging (MRI) is one of the most useful diagnostic tools in medicine today. The technique is based on the discovery that atomic nuclei possess a magnetic moment which can be detected using a magnetic field. When placed in a magnetic field, nuclei arrange themselves in a manner that aligns their magnetic moments either with or against the direction of the magnetic field. Those that align against the field (so-called β-spin nuclei) are higher in energy than those aligned with the field (so-called α-spin nuclei). The energy difference between α- and β-spin nuclei is directly proportional to the strength of the applied field. To measure this

difference in energy, the nuclei are subjected to electromagnetic radiation. Low energy α -spin nuclei absorb electromagnetic radiation causing them to "flip" against the field (or out of alignment with the field) 5 into a high energy B-spin. The frequency of the absorbed radiation, known as the "resonance frequency," reveals the α -B energy difference. Alternatively, high energy B-spin nuclei or non-aligned nuclei may "relax" into a low energy α -spin state with concomitant release 10 of electromagnetic radiation. Both the resonance frequency and the relaxation rates (measured in terms of relaxation rate $(1/T_1)$ wherein T_1 = the time that it takes for non-aligned nuclei to relax) for any nucleus is dependent on the magnetic environment surrounding 15 that nucleus. For example, nuclei within a substantial (strong) electron shell are shielded from external magnetic fields and as a result possess a smaller α -B energy difference, compared to nuclei within a weak electron shell. Also, nuclei situated in the proximity 20 of other paramagnetic nuclei relax faster and have higher relaxation rates than those for which such a relaxation mechanism is not available.

It is the difference in the nuclear magnetic relaxation rates of water protons that is most commonly used as the source of information in diagnostic MRI. Water is ubiquitously found in soft tissue. The hydrogen atoms in water possess paramagnetic nuclei (protons) that, when subjected to resonance frequency and relaxation-rate measurements, provide direct information regarding their physical microenvironment. The information can then be processed through computational techniques to obtain detailed anatomical images. Inspection of these images can reveal the presence of diseased or abnormal tissue, lesions and fractures or other malfunctions that may be present.

Not surprisingly, MR imaging of some tissue is more difficult than others. This may be due to any number of factors which result in low contrast and hence lack of resolution in the MR images obtained. 5 Previous efforts to overcome such problems have led to the discovery of so-called contrast enhancement agents. Typically, such agents contain a paramagnetic metal ion which is capable of altering the relaxation rates of water protons in its proximity. Upon administration to 10 a patient, the metal-containing contrast agent is absorbed by various organs depending on the patient's metabolic and excretion pathways. Once absorbed, the contrast agent alters the relaxation rates of water protons in the organ or tissue in which it resides. 15 The MR images of that organ or tissue thus achieve enhanced contrast with respect to neighboring tissue which contains lower concentrations or none of the

Because of the acute toxicity of most

20 paramagnetic metals, however, ordinary inorganic salts
of paramagnetic metals are unsatisfactory as contrast
agents. A solution to this problem is to use an
organic chelating ligand or metal-sequestering agent.
Through complexation to the metal, the organic

25 chelating ligand would prevent release of free, toxic
metal yet allow proton relaxation enhancement by acting
as a non-toxic paramagnetic carrier.

paramagnetic agent.

To be effective, MRI contrast agents
(paramagnetic metal-ligand complexes) must satisfy
30 several criteria. They must be stable and have high
formation constants so that release of the toxic metal
is prevented. They must be sufficiently soluble in
aqueous solutions to facilitate their administration to
a patient. And, they must be capable of efficient
35 enhancement of relaxation rates of water protons in

25

solution. Efficiency is generally measured in terms of "relaxivity", which is defined as the increase in relaxation rate per concentration of the paramagnetic complex measured in units of mM.

Gries et al. have described complexes for use 5 as diagnostic agents in United States patent 4,647,447. Also, the active paramagnetic ingredient of the FDAapproved MRI contrast agent Magnevist® is a complex of diethylenetriaminepentacetic acid and gadolinium (III). 10 In United States patent 4,899,755, Lauffer and Brady describe how paramagnetic metal-ligand complexes can be designed and synthesized to target specific tissue for MRI enhancement. This tissue-specific approach provides several improvements over the previously 15 reported non-specific methods. Qualitatively, tissuespecific agents provide better MR images of the targeted tissues such as the liver and the bile duct. Quantitatively, tissue-specific contrast agents can be used in lower concentrations to achieve image 20 enhancements similar to that observed with non-specific agents at higher doses. Thus, previously undetectable (or difficult to detect) liver tumors or malfunctions in the biliary system can be detected using hepatobiliary-specific contrast agents.

Another improvement upon the non-specific methods of the art is provided in United States patent 4,880,008 where the relaxivity-enhancement of contrast agents is shown to improve upon noncovalent binding to specific tissue proteins. This strategy not only 30 allows targeting of specific tissue containing the binding proteins, it also provides image enhancement at even lower doses than previously possible (due to improved relaxivity enhancement of the protein-bound agents).

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In spite of these advances, there remains a need for MRI contrast agents that combine favorable stability, solubility, relaxivity-enhancement and protein-binding characteristics with a tissue-specific pharmaco-kinetic profile.

SUMMARY OF THE INVENTION

The present invention provides improved compounds that are useful as tissue-specific MRI contrast agents, compositions containing those

10 compounds and methods for using them. In particular, the present invention provides paramagnetic metal complexes of 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid and pharmaceutically acceptable salts thereof as useful and efficient agents for improved tissue-specific enhancement of contrast in MRI of mammalian organs and tissues.

DETAILED DESCRIPTION OF THE INVENTION

The tissue-specific contrast agents of this invention comprise complexes of a paramagnetic metal 20 and 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid and pharmaceutically acceptable salts thereof. The stereochemistry at position 1 may be R or S, and the paramagnetic complex may be used in racemic or enantiomerically pure form. Preferred paramagnetic 25 metals of this invention are selected from the group consisting of Gd(III), Fe(III), Mn(II and III), Cr(III), Cu(II), Dy(III), Tb(III), Ho(III), Er(III) and Eu(III). The most preferred paramagnetic metal is Gd(III).

Pharmaceutically acceptable salts of this invention include those derived from inorganic or organic acids and bases. Included among such acid salts are the following: acetate, adipate, alginate,

aspartate, benzoate, benzenesulfonate, bisulfate, butyrate, citrate, camphorate, camphorsulfonate, cyclopentanepropionate, digluconate, dodecylsulfate, ethanesulfonate, fumarate, glucoheptanoate, 5 glycerophosphate, hemisulfate, heptanoate, hexanoate, hydrochloride, hydrobromide, hydroiodide, 2hydroxyethanesulfonate, lactate, maleate, methanesulfonate, 2-naphthalenesulfonate, nicotinate, oxalate, pamoate, pectinate, persulfate, 3-phenyl-10 propionate, picrate, pivalate, propionate, succinate, tartrate, thiocyanate, tosylate and undecanoate. Base salts include ammonium salts, alkali metal salts, such as sodium and potassium salts, alkaline earth metal salts, such as calcium, magnesium and zinc salts, salts 15 with organic bases, such as dicyclohexylamine salts, Nmethyl-D-glucamine, and salts with amino acids such as arginine, lysine, and so forth. Also, the basic nitrogen-containing groups can be quaternized with such agents as lower alkyl halides, such as methyl, ethyl, 20 propyl, and butyl chloride, bromides and iodides; dialkyl sulfates, such as dimethyl, diethyl, dibutyl and diamyl sulfates, long chain halides such as decyl, lauryl, myristyl and stearyl chlorides, bromides and iodides, aralkyl halides, such as benzyl and phenethyl 25 bromides and others. Water or oil-soluble or dispersible products are thereby obtained. The preferred salts of this invention are the N-methyl-Dglucamine, calcium and sodium salts.

The pharmaceutical compositions of this

invention comprise any of the complexes of the present invention, or pharmaceutically acceptable salts thereof, together with any pharmaceutically acceptable carrier, adjuvant or vehicle. Pharmaceutically acceptable carriers, adjuvants and vehicles that may be used in the pharmaceutical compositions of this

invention include, but are not limited to, ion
exchangers, alumina, aluminum stearate, lecithin, serum
proteins, such as human serum albumin, buffer
substances such as phosphates, glycine, sorbic acid,

5 potassium sorbate, TRIS (tris(hydroxymethyl)aminomethane), partial glyceride mixtures of saturated
vegetable fatty acids, water, salts or electrolytes,
such as protamine sulfate, disodium hydrogen phosphate,
potassium hydrogen phosphate, sodium chloride, zinc

10 salts, colloidal silica, magnesium trisilicate,
polyvinyl pyrrolidone, cellulose-based substances,
polyethylene glycol, sodium carboxymethylcellulose,
polyacrylates, waxes, polyethylene-polyoxypropyleneblock polymers, polyethylene glycol and wool fat.

According to this invention, the 15 pharmaceutical compositions may be in the form of a sterile injectable preparation, for example a sterile injectable aqueous or oleaginous suspension. This suspension may be formulated according to techniques 20 known in the art using suitable dispersing or wetting agents and suspending agents. The sterile injectable preparation may also be a sterile injectable solution or suspension in a non-toxic parenterally-acceptable diluent or solvent, for example as a solution in 1,3-25 butanediol. Among the acceptable vehicles and solvents that may be employed are water, Ringer's solution and isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending medium. For this purpose, any 30 bland fixed oil may be employed including synthetic mono- or di-glycerides. Fatty acids, such as oleic acid and its glyceride derivatives are useful in the preparation of injectables, as are natural pharmaceutically-acceptable oils, such as olive oil or 35 castor oil, especially in their polyoxyethylated

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versions. These oil solutions or suspensions may also contain a long-chain alcohol diluent or dispersant, such as Ph. Hely or similar alcohol.

The compounds and pharmaceutical compositions 5 of the present invention may be administered orally, parenterally, by inhalation spray, topically, rectally, nasally, buccally, vaginally or via an implanted reservoir in dosage formulations containing conventional non-toxic pharmaceutically-acceptable 10 carriers, adjuvants and vehicles. The term "parenteral" as used herein includes subcutaneous, intravenous, intramuscular, intra-articular, intrasynovial, intrasternal, intrathecal, intrahepatic, intralesional and intracranial injection or infusion 15 techniques.

When administered orally, the pharmaceutical compositions of this invention may be administered in any orally acceptable dosage form including, but not limited to, capsules, tablets, aqueous suspensions or 20 solutions. In the case of tablets for oral use, carriers which are commonly used include lactose and corn starch. Lubricating agents, such as magnesium stearate, are also typically added. For oral administration in a capsule form, useful diluents 25 include lactose and dried corn starch. When aqueous suspensions are required for oral use, the active ingredient is combined with emulsifying and suspending agents. If desired, certain sweetening, flavoring or coloring agents may also be added.

Alternatively, when administered in the form of suppositories for rectal administration, the pharmaceutical compositions of this invention may be prepared by mixing the agent with a suitable nonirritating excipient which is solid at room temperature 35 but liquid at rectal temperature and therefore will

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melt in the rectum to release the drug. Such materials include cocoa butter, beeswax and polyethylene glycols.

As noted before, the pharmaceutical compositions of this invention may also be administered topically, especially when the target of treatment includes areas or organs readily accessible by topical application, including the eye, the skin, or the lower intestinal tract. Suitable topical formulations are readily prepared for each of these areas or organs.

Topical application for the lower intestinal tract can be effected in a rectal suppository formulation (see above) or in a suitable enema formulation. Topically-transdermal patches may also be used.

For topical applications, the pharmaceutical 15 compositions may be formulated in a suitable ointment containing the active component suspended or dissolved in one or more carriers. Carriers for topical administration of the compounds of this invention include, but are not limited to, mineral oil, liquid petrolatum, white petrolatum, propylene glycol, polyoxyethylene, polycxypropylene compound, emulsifying wax and water. Alternatively, the pharmaceutical compositions can be formulated in a suitable lotion or 25 cream containing the active components suspended or dissolved in one or more pharmaceutically acceptable carriers. Suitable carriers include, but are not limited to, mineral oil, sorbitan monostearate, polysorbate 60, cetyl esters wax, cetearyl alcohol, 30 2-octyldodecanol, benzyl alcohol and water.

For ophthalmic use, the pharmaceutical compositions may be formulated as micronized suspensions in isotonic, pH adjusted sterile saline, or, preferably, as solutions in isotonic, pH adjusted sterile saline, either with our without a preservative

such as benzylalkonium chloride. Alternatively, for ophthalmic uses, the pharmaceutical compositions may be formulated in an ointment such as petrolatum.

For administration by nasal aerosol or
inhalation, the pharmaceutical compositions of this
invention are prepared according to techniques wellknown in the art of pharmaceutical formulation and may
be prepared as solutions in saline, employing benzyl
alcohol or other suitable preservatives, absorption
promoters to enhance bioavailability, fluorocarbons,
and/or other conventional solubilizing or dispersing
agents.

The amount of active ingredient that may be combined with the carrier materials to produce a single dosage form will vary depending upon the host treated, the sensitivity of the MR imaging instrument, the target tissue of the imaging experiment, the particular mode of administration and the intended effect of use. It should be understood, however, that a specific dosage regimen for any particular patient will depend upon a variety of factors, including the activity (induced relaxivity) of the specific agent employed, the age, body weight, general health, sex, diet, time of administration, rate of excretion, drug combination, and the judgment of the treating physician.

Dosage levels of between about 0.1 and about 1000 µmol/kg body weight per day, preferably between about 0.5 and about 300 µmol/kg body weight per day of the active ingredient compound are useful. A typical preparation will contain between about 1-1000 mM of the active complex. Preferably, such preparations contain between about 10-500 mM of the active complex.

To preserve shelf stability, to enhance the safety of the contrast agents and to prevent <u>in vivo</u>

35 release of toxic metal (or ligand), according to one

embodiment of this invention the compositions further contain a free organic ligand or an excess of 1-(p-nbutylbenzyl)diethylenetriamine pentaacetic acid and pharmaceutically acceptable salts of such ligands, for 5 example, calcium, sodium and meglumine (N-methyl-Dglucamine) salts and combination salts such as a calcium sodium salt. The pharmaceutically acceptable compositions contain between 0-5% (molar ratio) of excess organic ligand or a pharmaceutically acceptable 10 salt thereof, preferably 0-1%, and most preferably 0-0.5%. The use of compositions with an excess of a chelating ligand in diagnostic imaging has been disclosed in greater detail in International patent publication WO 90/03804, European patent publications 15 EP-A-0463644 and EP-A-454078 and United States patent 5,078,986.

The tissue-specific contrast agents of this invention may be readily obtained using conventional techniques. Scheme 1 illustrates a representative

20 example of a process for the preparation of gadolinium(III)-1-(p-n-butylbenzyl)diethylenetriamine pentaacetate dimeglumine salt (MS-264). It will be appreciated by those of skill in the art that the below synthetic processes are not intended as a comprehensive list of all means by which the contrast agents or the intermediates of this invention may be synthesized. Further methods or modifications of the below process will be evident to those of skill in the art.

starting with 4-n-butylaniline, the process
involves diazotization, followed by metal-catalyzed
coupling with acrylonitrile or an equivalent (e.g., an
alkyl ester of acrylic acid). The product is converted
to p-n-butylphenylalanine which is then processed to
obtain its ethylenediamine amide. Reduction of the
amide to the amine using a known reducing agent,

Scheme 1

$$\frac{\text{N-Methyl-D-Glucamine}}{0.5 \text{ Gd}_2\text{O}_3} \begin{bmatrix} \text{N} & \text{CO}_2 \\ \text{Bu} & \text{CO}_2 \\ \text{CO}_2 & \text{Gd}^{3+} & \text{CO}_2 - \end{bmatrix}^2 \cdot \begin{bmatrix} \text{CH}_2\text{NH}_2\text{CH}_3} \\ \text{H} & \text{OH} \\ \text{HO} & \text{H} \\ \text{H} & \text{OH} \\ \text{CH}_2\text{OH} \end{bmatrix}^{2+} \\ 10$$

preferably borane-THF complex, provides the p-n-butylbenzyl derivative of diethylenetraimine. The nitrogen atoms in this compound are alkylated using an alkyl ester of bromoacetic acid such as t-butyl-bromoacetate to give, after ester hydrolysis, the desired organic ligand.

Complexation to the paramagnetic metal can be achieved using any of the well-known procedures in the art. Scheme 1 provides an example of such a procedure involving the reaction of the ligand with gadolinium oxide in the presence of N-methyl-D-glucamine. Similar procedures using other paramagnetic metal oxides or salts instead of Gd₂O₃ and other bases instead of N-methylglucamine provide further paramagnetic contrast agents of this invention.

Optically active contrast agents may also be prepared using, for example, optically active p-n-butylphenylalanine or using other optically active intermediates in the process illustrated in Scheme 1.

20 Optically active intermediates can, in turn, come from resolution of a corresponding racemic mixture or from stereoselective synthesis. Methods for both resolution of racemic mixtures and stereoselective synthesis of optically active compounds are well known and require no further description here.

The contrast agents of this invention are surprisingly efficient and tissue-specific in MRI of mammalian tissue. Thus, this invention provides a method for tissue-specific contrast enhancement of MR images of mammalian organs and tissues. In particular, the method allows for contrast enhancement of MR images of the hepatobiliary system. For example, MS-264, the preferred contrast agent of this invention, exhibits surprisingly superior liver and bile specificity relative to structurally similar contrast agents

previously disclosed (see Examples section). MS-264 can also be used for tissue-specific imaging of tumors and the blood pool.

Following administration of the contrast 5 agent, NMR imaging is carried out; the choice of pulse sequence (inversion recovery, IR; spin echo, SE; gradient echo, GE; turbo FLASH,; etc.) and the values of the imaging parameters (echo time, TE; inversion time, TI; repetition time, TR; flip angel, etc.) will 10 be governed by the diagnostic information sought. In general, if one desires to measure T_1 , then TE should be less than 30 milliseconds (or the minimum value) to maximize T,-weighting. Conversely, if one desires to measure T_2 , then TE should be greater than 30 15 milliseconds to minimize competing T₁ effects. TR will remain approximately the same for both T_1 - and T2-weighted images; TI and TR are generally on the order of about 200-600 and 100-1000 milliseconds, respectively.

It is well known that ligands form complexes that are useful as MRI contrast agents may also be used in chelate complexes for diagnostic X-ray imaging. Therefore, according to another embodiment of this invention, metal complexes of 1-(p-n-butylbenzyl) diethylenetriamine pentaacetic acid and pharmaceutically acceptable salts thereof are used for diagnostic X-ray imaging.

In order that this invention may be more fully understood, the following examples are set forth.

These examples are for the purpose of illustration only and are not to be construed as limiting the scope of the invention in any way.

EXAMPLES

Relaxivity in rat liver cytosol

Rat liver cytosol was prepared by homogenizing rat livers (25% w/v) in 0.25 M sucrose/0.01 M 5 phosphate buffer, pH 7.4, centrifuging at 100,000 g for 90 min at 4°C, and concentrating the supernatant by pressure ultrafiltration to get the cytosol protein concentration close to that in vivo (5-10%). relaxivity of the Gd(III) complexes in cytosol was 10 measured at 37 °C using a Bruker 20 MHz Minispec equipped with a variable temperature probe. inversion-recovery pulse sequence with 10 data points was used for T_1 measurements. $1/T_1$ relaxation rates of solutions with varying concentrations of the chelates 15 (0.01-0.5 mM) were determined. The limiting slope of the $1/T_1$ vs concentration plot at low concentrations of the chelates in units of s⁻¹ mM⁻¹ was taken as the relaxivity (R1) of the chelate in cytosol. summarized in TABLE 1 (Bn = benzyl).

20 Higher relaxivity values imply higher binding affinity to cytosolic proteins and/or greater relaxivity when bound.



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TABLE 1. Relaxivity In Rat Liver Cytosol

	DTPA derivative	R1 in liver cytosol 20 MHz 37°C (s mM)
5	DTPA (R=R'=H) BOPTA (R=H, R'=BnOCH ₂)	7.1 11.3
10	R (R'=H) 1-(benzyl) 2-(benzyl) 1-(phenylethyl) [CH ₂ CH ₂ Ph]	11.7 nd 13.6
	1-(p-ethoxybenzyl) [EOB-DTPA] 1-(p-ethylbenzyl) 1-(p-isopropylbenzyl) 1-(p-n-butylbenzyl) [MS-264]	14.4 16.4 18.7 23. 8
15	1-(2,4-di-Me-benzyl) 1-(2,4,6-tri-Me-benzyl)	15.3 17.9
	1,3-dibenzyl 1,4-dibenzyl	23.7 20.0
20	1-(p-ethylbenzyl)-3-benzyl 1,3-di(p-ethylbenzyl) 1-(p-n-butylbenzyl)-3-benzyl	nd nd nd

Rat liver uptake and 1/T1 relaxation rates of rat liver

Un-anesthetized adult male Sprague Dawley rats (200-300 g) were injected in the tail vein with 50 µmol/kg of gadolinium chelate labeled with ¹⁵³Gd.

Rats (approx. 3 per time period) were euthanized by cervical dislocation (while anesthetized by CO₂) at 1, 5 and 30 min. post-injection. The liver was analyzed for ¹⁵³Gd concentration using a gamma counter. The percentage of the injected dose (% ID) in the liver was determined for each time point and the integrated area under the mean liver % ID time curve (AUC) was calculated.

For the 5 min. time period, the 1/T₁ proton relaxation rate of the excised liver was measured at 20 MHz and 37 °C as described above. Higher 1/T₁ values are associated with greater MRI signal intensity on T₁-weighted images and are due to greater liver uptake of the agent and/or greater relaxivity in the microenvironments of the liver. Results are summarized in TABLE 2.

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TABLE 2. Uptake and Induced Proton Relaxation Rates in Rat Liver

5	DTPA derivative	Area under the % inj. dose in rat liver vs time curve (AUC) from 0 to 30 min*	1/T1 of excised rat liver 5 min. post-inj. (s ⁻¹ ; 20 MHz, 37°C)
	DTPA (R=R'=H) BOPTA (R=H, R'=BnOCH ₂)	35 (n = 9) nd	4.64 ± 0.16 (n-3)
10	R (R' = H) 1-(benzyl) 2-(benzyl) 1-(phenylethyl) [CH2CH2Ph]	213 (n=9) nd nd	nd nd nd
15	1-(p-ethoxybenzyl) [EOB-DTPA]	408 (n=9)	10.1 ± 1.5 (n=3)
	1-(p-ethylbenzyl)	469 (n=10)	12.3 ± 0.9 (n=3)
	1-(p-isopropylbenzyl)	400 (n=8)	10.9 ± 1.2 (n=3)
	1-(p-n-butylbenzyl) [MS-264]	491 (n=6)	14.6 ± 0.2 (n=2)
	1-(2,4-di-Me-benzyl)	404 (n = 7)	10.2 ± 0.6 (n=3)
	1-(2,4,6-tri-Me-benzyl)	408 (n = 7)	11.7 ± 0.6 (n=3)
	1,3-dibenzyl	254 (n = 9)	8.40 (n = 1)
	1,4-dibenzyl	378 (n = 3)	8.66 (n = 1)
20	1-(p-ethylbenzyl)-3-benzyl	nd	nd
	1,3-di(p-ethylbenzyl)	260 (n=3)	11.2 (n = 1)
	1-(p-n-butylbenzyl)-3-benzyl	313 (n=3)	13.0 ₁ (n = 1)

 ⁽Total number of animals in experiment listed as "n")

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Rabbit liver uptake

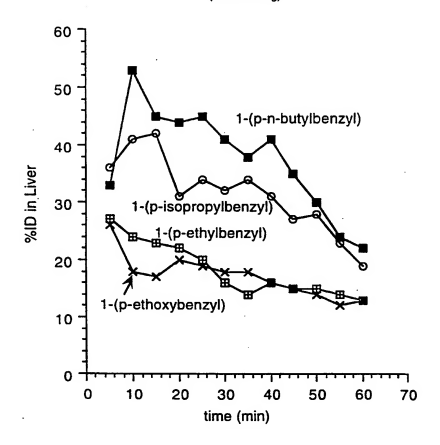
Anesthetized (pentobarbital, 35 mg/kg, ip.) adult rabbits (2.5-4 kg) were administered 50 \(\mu\text{mol/kg}\) of a 153 Gd-labeled chelate (approximately 30 μ Ci/kg) via 5 a cannulated marginal ear vein. Rabbits (1 per compound) were positioned supine under a gamma camera set with a photo peak energy of 100 keV +/- 20% and equipped with a low energy parallel hole collimator. Immediately after chelate administration, dynamic 10 planar whole body scintigraphy was performed every five minutes for at least one hour. Images were acquired in a 64 x 64 matrix, and region of interest (ROI) analysis was performed using the Image 1.52 software package (National Institutes of Health). Percent injected does 15 (% ID) in the liver at 5 min. intervals was calculated by dividing the counts in the liver ROI by the total counts in the body. Results are summarized in TABLE 3 and GRAPH 1.

TABLE 3

20	$ \begin{bmatrix} R & CO_2 \\ 1 & N & 4 & CO_2 \\ N & N & CO_2 \end{bmatrix} \cdot Gd(III) $ $ R = R CO_2 $	Area under the % inj. dose in rabbit liver vs time curve (AUC) 0 to from 30 min (n=1 rabbit per cmpd)
	1-(p-ethoxybenzyl) [EOB-DTPA]	480
	1-(p-ethylbenzyl)	553
	1-(p-isopropylbenzyl)	910
	1-(p-n-butylbenzyl) [MS-264]	1120

GRAPH 1

Rabbit Liver Scintigraphy of Gd153 Complexes (0.05mmol/kg)



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Monkey liver uptake

Two adult male cynomolgus monkeys were scintigraphically imaged after iv. administration of 50 μmol/kg of the chelates labeled with ¹⁵³Gd
5 (approximately 40 μCi/kg). All agents were given to each monkey on separate days. Monkeys were anesthetized with ketamine hydrochloride (i.m., to effect), intubated and maintained on isoflurane (0.5-2%) and positioned supine under a gamma camera prior to compound administration. Images were acquired and analyzed as described above. Results are summarized in TABLE 4 and GRAPHS 2 and 3.

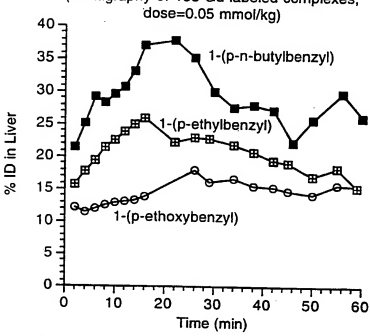
TABLE 4

15	$\begin{bmatrix} & & & & & & & & & & & & & & & & & & &$	Area under t % inj. dose liver vs tim (AUC) from (in monkey
	R	Monkey # 1	Monkey #2
20	1-(p-ethoxybenzyl) [EOB-DTPA] 1-(p-ethylbenzyl) 1-(p-n-butylbenzyl) [MS-264]	396 617 912	380 494 643

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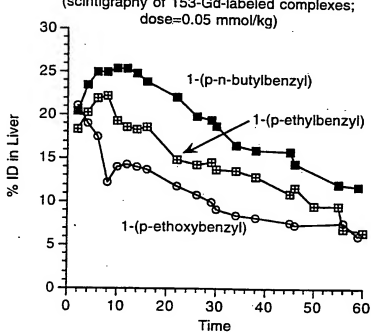
GRAPH 2

Liver Uptake in Monkey #1 (scintigraphy of 153-Gd-labeled complexes;



GRAPH 3

Liver Uptake in Monkey #2 (scintigraphy of 153-Gd-labeled complexes;



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Biliary excretion in monkeys

Two adult male cynomolgus monkeys used for scintigraphic imaging were injected on different days with iv bolus of 50 μ mol/kg of the chelates labeled 5 with 153 Gd (approximately 40 μ Ci/kg). Following completion of the imaging study, the animals were placed in metabolic cages and their total urine void and excreted feces were measured using a gamma counter at 24 and 48 hr post compound administration. The 10 average recovery of activity was 68% of the injected dose. Results are summarized in TABLE 5.

TABLE 5.

15	$\begin{bmatrix} & & & & & & & & & & & & & & & & & & &$	<pre>% of recovered activity in feces in (biliary excretion) of monkeys over 48 hr (n=2)</pre>
20	1-(p-ethoxybenzyl) [EOB-DTPA] 1-(p-ethylbenzyl) 1-(p-n-butylbenzyl) [MS-264]	35 ± 3% 48 ± 10% 86 ± 5%



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Synthesis of MS-264

α-Chloro-β-p-n-butylphenylpropionitrile (2)

One and a half moles (237 mL) of 4-nbutylaniline was dissolved in acetone (1500 mL). 5 solution was cooled to 10 °C and water (150 mL) and concentrated aqueous HCl (300 mL) were added. Sodium nitrite (69.0 g, 1.73 moles) in water (200 mL) was added dropwise to the cooled solution. The solution was adjusted to pH 4 by addition of solid sodium 10 acetate. Acrylonitrile (148 mL, 2.25 moles) was added, followed by $CuCl_2.2H_2O$ (38.4 g) in 50 mL H_2O . reaction mixture was heated to 40°C and was stirred at this temperature for 3 h. The aqueous layer was extracted with ether. The combined organics were 15 washed with water and saturated aqueous NaHCO3, were dried over Na2SO4, filtered and concentrated under reduced pressure. The residue was fractionally distilled at reduced pressure. The product was collected at 115-130°C @ 2 mm Hg, as 166.3 g (50% 20 yield) of an orange oil.

α-Chloro-β-p-n-butylphenylpropionic Acid (3)

The α-chloro-β-p-butylphenylpropionitrile 2

(166 g, 0.750 moles) was stirred over 240 mL formic acid and 160 mL concentrated HCl at reflux for 16 h.

25 The reaction mixture was cooled, diluted with 300 mL water and washed with ethyl acetate. The ethyl acetate solution was washed with water, saturated aqueous NaCl, dried over Na₂SO₄, filtered and concentrated under reduced pressure. The bulk oil solidified on standing.

30 The solid was recrystallized twice from hexane, yielding 139.0 g (0.58 moles, 77% yield) of a white solid, mp. 52.7-53.6°C.

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p-n-Butylphenylalanine (4)

α-Chloro-β-p-butylphenylpropionic acid 3

(128 g, 0.534 moles) was suspended in 800 mL 30% aqueous ammonia. The reaction was stirred for 4 days at 40°C in a tightly stoppered flask. After cooling to room temperature, the precipitate was removed by filtration through a coarse glass frit. The solid was washed thoroughly with ethanol followed by diethyl ether. The solid was dried under vacuum yielding 54.3 g (0.245 moles, 46% yield) of p-butylphenylalanine as a white solid. The solid was recrystallized from 2N HCl to give a white solid, mp 218-223°C.

p-n-Butylphenylalanine Methyl Ester Hydrochloride (5)

Butylphenylalanine 4 (4.20 g, 19.0 mmoles)

was suspended in 50 mL dry methanol. The solution was saturated with HCl gas and was refluxed for 2 hours. The solution was cooled and then added to 400 mL diethyl ether. The resulting solid was collected by vacuum filtration and was washed with ether. The solid was redissolved in a minimum volume of methanol and precipitated in ether as before resulting in 3.45 g (12.7 mmoles, 67% yield) of white solid (mp 164.5-165.1°C).

p-n-Butylphenylalanine Ethylenediamine Amide (6)

Butylphenylalanine methyl ester hydrochloride
5 (3.45 g, 12.7 mmoles) was dissolved in 65 mL
ethylenediamine and was stirred at room temperature for
16 h. The ethylenediamine was removed by evaporation
at reduced pressure. The residue was partitioned

between 1 N NaOH and ethyl acetate and the aqueous was
washed three times with ethyl acetate. The combined
organic solutions were dried over Na₂SO₄, filtered and
concentrated yielding 2.76 orange oil (83% yield). ¹H



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NMR (CDCL₃) & 7.50 (br. s, 1H), 7.15 (s, 4H), 3.60 (dd, 1H), 3.38-3.17 (3H), 2.85-250 (5H), 1.58 (m, 2H), 1.42-1.24 (6H), 0.93 (t, 3H).

1-p-n-Butylbenzyldiethylenetriamine Trihydrochloride (7)

Butylphenylalanine ethylenediamine amid 6 (2.76 g, 10.5 mmoles) was dissolved in 50 mL THF. Borane THF (50 mL, 1.0 M) was added slowly to the stirred solution. The reaction was then refluxed under argon for 16 h, and quenched by careful addition of methanol. The reaction mixture was concentrated under reduced pressure. Dry ethanol (75 mL) was added and the solution was saturated with HCl gas at 0 °C. The mixture was brought to reflux for 24 h, after which the 15 solution was cooled and concentrated. Ethyl ether was added to the concentrated solution and the resulting precipitate was collected by vacuum filtration and washed with ether, yielding 3.45 g (9.6 mmoles, 92%) white solid. Analysis calc'd for C₁₅H₂₇N₃•3HCl: C, 20 50.22; H, 8.43; N, 11.71; Cl, 29.64 found: C, 50.16; H, 8.44; N, 11.76; Cl, 29.57.

<u>1-p-n-Butylbenzyldiethylenetriamine penta-t-butyl</u> Acetate (8)

Butylbenzyldiethylenetriamine

trihydrochloride 7 (3.45 g, 9.62 mmoles) was suspended in 50 mL dry DMF. Diisopropylethylamine (25 mL, 144 mmoles) was added followed by 12.5 mL (77 mmoles) to butylbromoacetate. The reaction was stirred at room temperature for 16 h under argon. The DMF and excess reagents were removed by evaporation under reduced pressure. The residue was partitioned between concentrated aqueous NaHCO3 and chloroform. The organic solution was washed once each with water and saturated aqueous NaCl, dried over Na2SO4, filtered and

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concentrated under reduced pressure. The crude product was filtered through silica gel in 1:1 ethyl acetate:hexane. After being reconcentrated, the product was obtained as 6.88 g (8.39 mmol, 87% yield) of a light yellow oil. ¹H NMR (CDCl₃) & 7.15 (d, 2H), 7.04 (d, 2H), 3.50-3.35 (10H), 3.09 (m, 1H), 2.83 (m, 2H), 2.75-245 (8H), 1.65-1.10 (49H), 0.94 (t, 3H).

1-p-n-Butylbenzyldiethylenetriamine Pentaacetic acid (9)

The penta-t-butyl ester 8 (6.88 g, 8.39 mmol) was dissolved in 1:1 dioxane:concentrated HCl and was stirred at room temperature for 16 h. The reaction mixture was concentrated under reduced pressure yielding the crude product as 4.79 g of a tan foam.

The product was eluted from a 10 g C-18 Sep-Pak cartridge with 40% aqueous acetonitrile. The acetonitrile was evaporated and the resulting aqueous solution was lyophilized producing 3.30 g (6.12 mmoles, 73% yield) of a flocculent white solid. Analysis calc'd for C25H37N3O10: C, 55.65; H, 6.91; N, 7.79; found: C, 55.06; H, 6.99; N, 7.71; Cl, 0.00.

Gadolinium(III) 1-(p-n-butylbenzyl)diethylenetriamine pentaacetate dimeglumine salt (10)

butylbenzyl)diethylenetriamine pentaacetic acid 9
(6.00 g, 10.0 mmol [90% by weight]) and N-methyl-Dglucamine (3.90 g, 20 mmol) were weighed in a 100-mL
round-bottom flask and 30 mL of distilled water was
added to it. The mixture was stirred at 95°C for six
hours. The resulting cloudy solution was loaded on a
10-g C-18 Sep-Pak® cartridge and the complex was eluted
with water. The solvent was removed on a rotary
evaporator, the resulting white solid residue was dried
in high vacuo for 24 hours at 45°C. Yield: 9.78 g



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(88 %). Anal. Calcd for $C_{39}H_{70}GdN_5O_{21}$: C, 42.50; H, 6.40; N, 6.35; Gd, 14.27. Found: C, 42.11; H, 6.08; N, 6.35; Gd, 14.03.

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CLAIMS:

- 1. A complex of a paramagnetic metal and 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid and pharmaceutically acceptable salts thereof.
- The complex according to claim 1, wherein the paramagnetic metal is selected from the group consisting of Gd(III), Fe(III), Mn(II), Mn(III), Cr(III), Cu(II), Dy(III), Tb(III), Ho(III), Er(III) and Eu(III).
- 3. The complex according to claim 1, wherein the paramagnetic metal is Gd(III).
 - 4. Gadolinium(III)-1-(p-n-butylbenzyl) diethylenetriamine pentaacetate dimeglumine salt.
- 5. The complex according to claim 1 wherein 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid is present in racemic form.
- 6. The complex according to claim 1 wherein 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid is present enantiomerically enriched or
 20 enantiomerically pure form.
 - 7. A pharmaceutical composition comprising a compound according to any one of claims 1 to 6 together with a pharmaceutically acceptable carrier, adjuvant or vehicle.
- 25 8. The pharmaceutical composition according to claim 7, further comprising a free organic ligand or a pharmaceutically acceptable salt thereof.

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- 9. The pharmaceutical composition according to claim 7, further comprising a free organic ligand or calcium, sodium, meglumine or combination salts thereof.
- 5 10. The pharmaceutical composition according to claim 8 or claim 9, wherein the organic ligand is 1-(p-n-butylbenzyl)diethylenetriamine pentaacetic acid.
- 11. A method for tissue-specific contrast enhancement of MR images of mammalian organs and tissues comprising the step of administering a diagnostically effective amount of a complex according to any one of claims 1 to 6.
- 12. The method according to claim 11 for contrast enhancement of MR images of the hepatobiliary15 system.
 - 13. Use of a compound according to any one of claims 1 to 6 for the manufacture of a MRI contrast agent.
- 14. A method for tissue-specific diagnostic
 20 X-ray imaging of mammalian organs and tissues
 comprising the step of administering a diagnostically
 effective amount of a complex according to any one of
 claims 1 to 6.

INTERNATIONAL SEARCH REPORT

International Application No. 95/04045

A. CLA_SII	FICATION DEJECT MATTER 51 K 49/00, A 61 K 49/04		
According to	International Patent Classification (IPC) or to both national classifica	ation and IPC	
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Minimum do	ocumentation searched (classification system followed by classification	symbols)	
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Documentati	ion searched other than minimum documentation to the extent that suc	th documents are included in the fields se	arched
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Electronic d	ata base consulted during the international search (name of data base	and, where practical, search terms used)	
C. DOCUM	ENTS CONSIDERED TO BE RELEVANT		Delement to claim No
Category *	Citation of document, with indication, where appropriate, of the rele	evant passages	Relevant to claim No.
A	EP, A, 0 405 704 (SCHERING AKTIENGESELI 02 January 1991 (02.01 abstract; claims.		1-14
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Fur	ther documents are listed in the continuation of box C.	Patent family members are listed	i in annex.
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"L" docum	nent which may throw doubts on priority claim(s) or	involve an inventive step when the	TOCALLICUL IZ COCCU STORIC
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	NL - 2280 HV Rijswijk Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl, Fax (+ 31-70) 340-3016	SCHNASS e.h.	

INTERDONAL SEARCH REPORT

International application No. CT/00 95/04045

Box I Observations with certain claims were found unsearchable (Continuation of item I of first sheet)
This international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X Claims Nos.: 11,12,14 because they relate to subject matter.
because they relate to subject matter not required to be searched by this Authority, namely:
Remark: Although claims 11,12,14 are directed to a method of treatment of the human and/or animal body by therapy (Rule 39.1(iv)PCT)the search was carried out and based on the alleged effects of the composition.
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3. Claims Nos.:
because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(2).
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
This International Searching Authority found multiple inventions in this international application, as follows:
As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2. As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
3. As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:
4. No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark on Protest The additional search fees were accompanied by the applicant's protest.
No protest accompanied the payment of additional search fees.
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ANHANG

zum internationalen Recherchenbericht über die internationale Patentanmeldung Nr.

ANNEX

to the International Search Report to the International Patent Application No. ANNEXE
au rapport de recherche international relatif à la demande de brevet
international n°

PCT/US 95/04045 SAE 108350

In diesem Anhang sind die Mitglieder der Patentfamilien der im obengenannten internationalen Recherchenbericht angeführten Patentdokumente angegeben. Diese Angaben dienen nur zur Unterrichtung und erfolgen ohne Gewähr.

This Annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The Office is in no way liable for these particulars which are given merely for the purpose of information.

La presente annexe indique les membres de la famille de brevets relatifs aux documents de brevets cités dans le rapport de recherche international visée ci-dessus. Les reseignements fournis sont donnés à titre indicatif et n'engagent pas la responsibilité de l'Office.

Im Recherchenbericht angeführtes Patentdokument Patent document cited in search report Document de brevet cité dans le rapport de recherche	Datum der Veröffentlichung Publication date Date de publication	Mitglied(er) der Paientfamilie Patent family member(s) Membre(s) de la famille de brevets	Datum der Veröffentlichung Publication date Date de publication	
EP A2 405704	02-01-91	4901244665744435028887555511942027080017942015229442 12472170800194201522942 12472170820977712062015229425 12420275556344045222445 124242651403140992 1299255 1299255 120022445 120022445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 12002445 1200244 1200244 1200244 1200244 1200244 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 120024 1200	941301311012999999110131111111111111111111	